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(54) Dispersant-viscosity improvers for lubricating oil compositions

Dispergier- und Viskositätsverbesserer für Schmierötzusammensetzungen Additifs pour compositions d'huile lubrifiante améliorant la dispersibilité et la viscosité

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Description

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[0001] This invention relates to dispersant-viscosity improvers for lubricating oils, and oil compositions and concentrates containing such dispersant-viscosity improvers.

BACKGROUND OF THE INVENTION

[0002] The viscosity of oils of lubricating viscosity is generally dependent upon temperature. As the temperature of the oil is increased, the viscosity usually decreases, and as the temperature is enduced, the viscosity usually increases.

[0003] The function of a viscosity improver is to reduce the extent of the decrease in viscosity as the temperature is created to the created or to reduce the extent of the increase in viscosity as the Interpretature is lowered, or both. Thus, a viscosity improver ameliorates the change of viscosity of an oil containing it with changes in temperature. The fluidity characteristics of the oil are improved.

[0004] Viscosily improvers are usually polymeric materials and are often referred to as viscosity index improvers. § [0005] Ester group containing polymers are well-known additives for improving the fulldiply characteristic of luthicating oils. Polyacrylate, particularly polymerhacrylate seter polymers are well-known and are widely used for this purpose. [0006] Dispersants are also well-known in the luthicating and Lispersants are employed in luthicatents to keep impurrities, particularly those formed during operation of machinery, in suspension rather than allowing them to deposit on the surfaces of luthicated posts.

[0007] Multifunctional additives that provide both viscosity improving properties and dispersant properties are likewise known in he art. Such products are described in numerous publications including plater klamenn, Lubricants and Related Products", Verlag Chemie Gmbh (1984), pp 185-193; C. V. Smalheer and R. K. Smith "Lubricant Additives", Lezius-Hiles Co. (1987), M.-W. Ranney, "Lubricant Additives", Noyes Data Corp. (1973), pp 92-145, M.-W. Ranney, "Lubricant Additives, Recent Developments", Noyes Data Corp (1979), pp 138-164; and M.-W. Ranney, "Synthetic Oils and Additives for Lubricants", Noyes Data Corp. (1980), pp 96-168. Each of these publications is hereby expressly incorporated herein by reference.

[0008] Derivatives of polyacrylate esters are well-known as additives for lubricants that provide not only improved viscosity characteristics, but also enhance dispersant properties of lubricants.

[0009] It is desirable that the viscosity improver or dispersant viscosity improver not adversely affect the low-temperature viscosity of the lubficant containing same. Frequently, while viscosity improvers or dispersant viscosity improvers enhance the high temperature viscosity characteristics of lubricating oil, that is, they reduce the loss of viscosity with increasing temperature, low temperature properties of the treated fubricant become worse.

[0010] One of the major requirements for automatic transmission fluids has been improved tow temperature performance as demonstrated by a maximum Brookfield viscosity of 20,000 centipols as 4-0°C. The viscosity modifier, which can comprise nearly 50 weight percent of the total additive system employed in an automatic transmission fluid can have a major impact on the low temperature performance. Such characteristics are also desirable in other applications such as in gear lubricants. The copolymers of this invention are also useful in many other lubricating oil compositions including, but not limited to engine oils, hydraulic oils, industrial oils, etc.

[0011] Accordingly, it is desirable to provide compositions that reduce the extent of loss of viscosity at high temperatures while not adversely increasing the low temperature viscosity of lubricating oil compositions.

[0012] We have now found it possible to provide: novel multi-purpose abfrant additives; multi-purpose additives directed to improving the viscosity and dispersant properties of a business improved in the interest or preparing such multi-purpose additives, additive constrates containing the novel multi-purpose lubricant additives of this invention; and lubricant having improved dispersant and viscosity properties.

SUMMARY OF THE INVENTION

[0013] The present invention is directed to a polyacrylate ester, and more particularly a polymethacrylate ester based dispersant-viscosity improver for lubricating oil compositions. Specifically, the dispersant viscosity improvers of this invention are nitrogen-containing copolymers having a number average molecular weight in the range from 10,000 to 300,000 and polydispersity values in the range from 1.5 to 5 comprising units derived from

(A) 30% to 50% by weight of alkyl acrylate ester monomers containing from 1 to 11 carbon atoms in the alkyl group; (B) 40 % to 70 % by weight of alkyl acrylate ester monomers containing from 12 to 24 carbon atoms in the alkyl group; and

(C) 0.5 % to 5 % by weight of at least one nitrogen containing monomer selected from vinyl substituted nitrogen haterocyclic monomers, dialkylaminoalkyl acrylate monomers, dialkylaminoalkyl acrylamide monomers and tertiary-alkyl acrylamides, provided that the total equals 100%.

[0014] The present invention also relates to additive concentrates and lubricating oil compositions containing the compositions of this invention and processes for preparing the compositions.

[0015] As used herein, the terms "acrylate" and "acrylamide" include the alkyl acrylates and alkyl acrylamides, and especially, methacrylates and methacrylamides.

5 [0016] Various preferred features and embodiments of the invention will be hereinafter described by way of non-limiting illustration.

[0017] According to the present invention a composition of matter suitable for use as a dispersant-viscosity improver for lubricating of compositions comprises nitrogen-containing copolymers derived from a mixture of alkyl acrylate ester monomers containing, on one hand, from 1 to 12 data to a little alkyl group and on the other hand from 12 to 24 carbon atoms in the alkyl group and at least one nitrogen-containing monomer as described in greater detail hereinbelow.

[0018] As used herein, the terms "hydrocarbon", "hydrocarbyl" or "hydrocarbon based" mean that the group being described has predominantly hydrocarbon character within the context of this invention. These include groups that are purely hydrocarbon in nature, that is, they contain only carbon and hydrogen. They may also include groups containing substituents or atoms which do not alter the predominantly hydrocarbon character of the group. Such substituents may include halo-, alkovy-, nitro-, etc. These groups also may contain hetero atoms. Suitable hetero atoms will be apparent to those skilled in the art and include, for example, sulfur, nitrogen and oxygen. Therefore, while remaining predominantly hydrocarbon in character within the context of this invention, these groups may contain atoms other than carbon present in a chain or ring otherwise composed of carbon atoms.

20 [0019] In general, no more than about three non-hydrocarbon substituents or hetero atoms, and preferably no more than one, will be present for every 10 carbon atoms in the hydrocarbon or hydrocarbon based groups. Most preferably, the groups are purely hydrocarbon in acture, that is they are essentially free of atoms other than carbon and hydrogen. [0020] Throughout the specification and claims the expression oil soluble or dispersible is used. By oil soluble abgresible is meant that an amount needed to provide the desired level of activity or performance can be incorporated by being dissolved, dispersed or suspended in an oil of fluriorating viscosity. Usually, this means that at least 0.001% by weight of the material can be incorporated in a lubricating oil composition. For a further discussion of the terms oil soluble and dispersible, as pricularly "stably bigersible", see U.S. Patent 4,320,019.

[0021] In the context of this invention the term 'copolymer' means a polymer derived from two or more different monomers. Thus, a polymer derived from mixture of, for example, methyl-, buthyl-, Cs.,1-r, and C2.94; exciptates is a copolymer as defined herein. The copolymers of this invention also contain units derived from nitrogen-containing monomers.

[0022] The nitrogen-containing copolymers of this invention may be prepared by several different processes. In one embodiment, the nitrogen-containing copolymer is obtained by reacting, together, the acrylate ester monomers and the nitrogen-containing monomer. In another embodiment, the nitrogen-containing monomer is grafted onto a preformed acrylate copolymer backbone.

[0023] The nitrogen-containing copolymers of this invention are frequently prepared in an organic diluent. It has been determined that specific amounts and types of diluent present in the integen-containing copolymers of the invention can have a pronounced effect on the low temperature viscosity properties of bloricants containing them.

40 The Alkyl Acrylate Ester Monomers

[0024] As stated hereinabove, the nitrogen-containing copolymer comprises units derived from (A) from 30 % to 60 % by weight of alkyl acrylate ester monomers containing from 1 to 11 carbon atoms in the alkyl group and (B) from 40% to 70 % by weight of alkyl acrylate ester monomers containing from 12 to 24 carbon atoms in the alkyl group. In an especially preferred embodiment, the alkyl acrylate ester monomers comprise alkyl methacrylate esters.

(025) The acrylate ester monomers can be prepared by conventional methods well-known to those of skill in the art. For example, acrylate ester monomers are most often prepared via the propylene oxidation process, a two stage vapor phase oxidation of propylene to acrylic acid, which is then esterified to the desired ester. Previously, the manufacture of acrylates involved the petrochemistry of materials such as acetylene, acrylonitrile and others.

[0026] For methacrylates, processes used often vary, depending on the desired monomer. The acetone cyanohydrin process involves the reaction of acotone with HCN to form acetone cyanohydrin which is then reacted with the desired alcohol to form the ester. Propylene carbonylation and many other processes are also used.

[0027] A variety of procedures are described in considerable detail in the section entitled "Acrylic and Methacrylic Ester Polymers" in the Encyclopedia of Polymer Science and Engineering, Vol. 1, pp. 247-251, Wiley-Interscience, New York (1985).

[0028] Many alkyl acrylate esters are commercially available. Suppliers include, Rohm and Haas; San Esters Corp, with offices in New York, New York; Mitsubishi Rayon Co. Ltd.; Polysciences, Inc., Warrington, Pennsylvania; Sartomer Co., Exton, Pennsylvania; and others.

The Nitrogen-Containing Monomer

[0029] The nitrogen-containing copolymers of this invention also comprise units derived from (C) from 0.5 % to 5 % by weight, and especially from 1.5 % to 2 5 % by weight of at least on enitrogen-containing monomer selected from the group consisting of vinyl substituted nitrogen heterocyclic monomers, dialkylaminoalkyl acrylate monomers, dialkylaminoalkyl acrylated and to start a containing and to start a

[0030] In one embodiment, the nitrogen-containing monomer is an N-vinyl substituted heterocyclic monomer. Examples of such monomers include N-vinyl imidazole, N-vinyl pyrrolidinone and N-vinyl caprolactam.

[0031] In another embodiment, the vinyl substituted heterocyclic monomer is vinyl pyridine.

[0032] In yet another embodiment, the nitrogen-containing monomer is a N,N-dialkylaminoalkyl acrylamide or acrylate wherein each alkyl or aminoalkyl group contains, independently, from 1 to about 8 carbon atoms.

[0033] In a further embodiment, the nitrogen-containing monomer is a tertiary-alkyl acrylamide, preferably tertiary butyl acrylamide.

The Diluent

[0034] As noted hereinabove, the copolymers of this invention may be prepared in the presence of a diluent. A diluent may also be added to a substantially diluent-free copolymer, usually by dissolving or dispersing the substantially diluent-free polymer in an appropriate diluent.

20 [0035] In one embodiment, the diluent is a mineral oil. In an especially preferred embodiment the mineral oil consists essentially of hydrotreated naphthenic oil.

[0036] The diluent may also be a synthetic oil. Common synthetic oils are ester type oils, polyolefin oilgomers or alkylated benzenes.

[0037] The diluent-containing copolymers of this invention are referred to herein as additive concentrates. Such as additive concentrates are then added, along with other desirable performance-improving additives, to an oil of lubricating viscosity to prepare the finished lubricant composition.

[0038] The additive concentrates preferably comprise from about 25% to about 90% by weight of copolymer, preferably from 35% to about 80% by weight, and from about 10% to about 75% by weight of diluent, preferably from about 20% to about 65 % by weight of diluent.

[0039] In one embodiment, selection of diluents having particular characteristics leads to enhanced performance of dispersant-viscosity improvers of this invention. For example, lubricating oil compositions comprising additive concentrates containing certain diluents together with the nitrogen-containing polymers of this invention have enhanced low temperature characteristics. Particularly valuable are lubricating oil compositions that display excellent viscosity characteristics at very low temperatures, for example, from 5°C to 4°C°C.

5 [0040] The certain diluents that Impart surprising and exceptional low temperature performance when used in conjunction with the dispersant viscosity improvers of this invention, have in common very low viscosity at very low temperatures. In particular they all display Brookfield viscosities (expressed in comptops) at -28°C ranging from about 50 to about 400, more preferably trom about 50 to about 400, more preferably trom about 50 to about 200. At -40°C useful oils have Prookfield viscosities (expressed in centipoise) ranging from about 100 to about 1050, more preferably from about 125 to about 500. Brookfield viscosities of are determined employing ASTM Procedure D-2983 described in greater datalt hereinafter. These particularly useful diluents display viscosities ASTM Procedure D-445) at 40°C rangins from about 2.5 to about 5.0 certislotes and at all clients display viscosities of SSTM Procedure D-445) at 40°C rangins from about 2.5 to about 5.0.

100°C ranging from about 1 to about 2.5 centistokes.
[0041] included among such useful dilluents are naphthenic oils, hydrotreated naphthenic oils, and alkylated aromatics, particularly alkylated benzense having at least one alkyl group containing from about 8 to about 24 carbon atoms, preferably from 12 to about 18 carbon atoms. Especially useful are hydrotreated naphthenic oils, examples being Risella G-07, Cross Oil Co's L-40, a 40 neutral hydrotreated naphthenic oil and L-60, which is a 80 neutral oil.

[0042] Low temperature viscosity (Broodkield Viscosity) of fluid lubricants is determined using ASTM Procedure 2983, Standard Test Method for Low Temperature Viscosity of Automotive Fluid Lubricants Measured by Brooklield Viscometer, which appears in the Annual Book of ASTM Standards, Section 5, ASTM, Philadelphia, PA, USA. This procedure employs a Brookfield Viscometer which is described in the procedure. The device is available from Brookfield Engineering Laboratories, Stoughton, MA, USA.

[0043] ASTM Procedure D-445 is described hereinafter.

The Nitrogen-Containing Copolymer

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[0044] The copolymers of this invention have a number average molecular weight (M_n) ranging from 10,000 to 300,000, more often from 20,000 to 150,000 frequently from 30,000 to 100,000.

[0045] Molecular weights of polymers are determined using well-known methods described in the literature. Exam-

ples of procedures for determining molecular weights are gel permeation chromatography (also known as size-exclusion chromatography) and vapor phase osmometry. These and other procedures are described in numerous publication

[0046] P.J. Flory, "Principles of Polymer Chemistry" Cornell University Press (1953), Chapter VII, pp 266-316, and [0047] "Macromolecules, an Introduction to Polymer Science", F. A. Bovey and F.H. Winslow, Editors, Academic Press (1979) pp 296-312.

[0048] W.W. Yau, J.J. Kirkland and D.D. Bly, "Modern Size Exclusion Liquid Chromatography", John Wiley and Sons, New York. 1979.

[0049] A measurement which is complementary to a polymer's molecular weight is the melt index (ASTM D-1238). Polymers of high melt index generally have low molecular weight and vice versa. Mooney Viscosity (ASTM Procedure D-1646-87) relates indirectly to polymer molecular weight. All other factors being equal, as molecular weight of the polymer incresses, so too does the Mooney viscosity.

[0050] ASTM Procedures D-1238 and D-1646-87 appear in Volumes 8 and 9, respectively, of the aforementioned Annual Book of ASTM Standards.

15 [0051] A preferred method, and the method employed to determine the molecular weights of polymers as set forth herein, is gel permeation chromatography (GPC) employing polyacrylate standards.

[0052] As noted hereinabove, the copolymers of this invention may be prepared by several different techniques. In one embodiment, the acrylate ester monomers and nitrogen-containing monomer are reacted together. In another embodiment, the acrylate esters are reacted to form an acrylate ester copolymer backbone onto which is grafted a nitrogen-containing monomer. In still another embodiment, a mixture of acrylate and nitrogen-containing monomers

may be grafted onto a preformed acrylate ester polymer backbone.

[0053] În the first embodiment, a mixture of monomers is charged to a reactor together with, if desired, diluterit and again, if desired, chair transfer agent. The materials are usually strived under a nitrogen atmosphere, diluterit and application initiator is conveniently added and the materials are heated to reaction temperature. The reaction is continued until the desired decree of powerbards in statisment.

[0054] In an alternative, and generally preferred embodiment, the monomers are polymerized incrementally. A mixture of monomers together with a polymerization initiator is prepared. A portion, typically 20% to 40 %, more often 33 %, of the mixture, is charged to a reactor with the balance being placed in an addition vessel. The reactants are heated under a nitrogen atmosphere until an exotinemic reaction is noted. When the exothermic reaction begins to subside, addition of the balance of the monmer-initiator muture is begun, while maintaining, whe heafting or cooling, as needed,

the desired reaction temperature.

[0055] In a particular embodiment of this incremental polymerization process, monomers (A) and (B) and the free radical initiator are first combined to form a first moture is whereupon from 20% to 40% of said first moture is mixed with at least 50%, preferably at least 70%, more often 100% of the intended charge of monomer (C) to form a second mixture. Said second mixture is then heated until an exotherm is noted and freaction is continued until the exotherm subsides. When the exotherm subsides, when the exotherm subsides, when the exotherm commoners (A) and (B) and initiator is begun while maintaining the desired reaction temperature until the addition is completed. The reaction is then continued to completion.

[0056] In the second embodiment, the acrylate monomers are polymerized, then the grafting of the nitrogen-containing monomer onto the preformed acrylate ester opolymer is accomplished. A mixture of additional acrylate monomers together with nitrogen-containing monomer may be grafted onto the preformed acrylate ester polymer.

[0057] Either the single step procedure or the incremental procedure outlined above may be used for polymerization of the acrylate monomers.

[0058] The entire charge of the nitrogen containing monomer may be present at the start of the polymerization process. Alternatively, the nitrogen containing monomer may be added to the already prepared polyacytiale either granully or incrementally. In the grafting process, additional initiator is usually employed during the grafting step. In either process. additional initiator may be added during processing.

[0059] Polymerization can take place under a variety of conditions, among which are bulk polymerization, solution polymerization, emulsion polymerization, suspension polymerization and nonaqueous dispersion techniques.

[0060] To prepare the copolymers constituting the dispersant viscosity improvers in accordance with the invention it is possible to make use of the conventional methods of radical copolymerization.

[0061] Such methods are described in the work "Encyclopedia of Polymer Science and Engineering" (H.F. Mark, N. M. Bikales, C.G. Overberger and G. Menges), 2nd edition (1988), published by Wiley Interscience.

[0062] These methods include free-radical initiated polymerization employing azo compounds or peroxides. Also described therein are photochemical and radiation initiated methods.

[0063] Useful initiators include organic peroxides, hydroperoxides and azo compounds.

[0064] Molecular weights of the polymers can be controlled employing a number of techniques including choice of initiator, reaction temperature, concentration of monomers and solvent type. Chain transfer agents can also be used.

[0065] Ionic polymerization techniques are known including cationic and anionic methods; however, cationic methods are generally ineffective for acrylate and methacrylate monomer polymerization.
[0066] Free radical initiation is preferred.

[0057] Free radical generating reagents useful as polymerization initiators are well known to those skilled in the art. Numerous examples of free radical initiators are mentioned in the above-referenced texts by Flory and Bovey and Winslow. An extensive listing of free radical initiators appears in J.Brandup and E.H. Immergut, Editors, "Polymer Handbook", 2nd edition, John Wiley and Sons, New York (1975), pp II-1 to II-40. Numerous free radical initiators are available and many are commercially available in large quantities. Included among free radical initiators are towly peroxide, I-buty hydroperoxide, I-army peroxide, curryl peroxide, dibenzoyl peroxide (Aldrich), I-butyI,m-chloroperbenzoale, azobisvalerohitile, I-butyl peroctade and iertiany-butyl perbenzoale, (Trigonox 21 and Trigonox C, respectively, both from ARXQ) and 2,2° azobis(cobutyronities), (VAZO-64, UPPont).

[0068] Free radical initiators are usually used in amounts ranging from 0.01% to 10 percent by weight based on the total weight of reaction mixture. Preferably, the initiators are used at 0.05% to 3% by weight, often up to 1% by weight. [0069] The choice of free radical initiator can be an important consideration. Considerations include the half-life of the initiator at a given temperature, nature of the reactants, reaction temperature, solvent or diluent, and the like.

[0070] The products of the present invention are generally prepared at temperatures ranging from 60°C to 140°C, more often from 80°C to 120°C, and preferably from 90°C to 110°C. The half-life of an initiator at a given temperature is an important consideration.

[0071] Because acrylic polymerizations are usually accompanied by liberation of considerable heat, care must be 20 taken to avoid uncontrolled reaction. Temperatures can be controlled by using reactors with cooling jackets, controlling rates of addition and use of reaction solvents.

[0072] Another useful means for preparing the copolymers of this invention is to employ a high energy mechanical mixing device. These include off limits, ball mills or extruders. Of these, extruders are preferred since the comonomes can be fed to the feed hopper in any desired fashion. Methods of employing such devices, and especially extruders, are described in a number of patients including havashi et al. U.S. 4870.173 and 5000 et al. U.S. 5.039.433.

[0073] The following examples are intended to illustrate several compositions of this invention as well as means for preparing same. Unless indicated otherwise all parts are parts by weight. It is to be understood that these examples are intended to illustrate several compositions and procedures of the invention and are not intended to limit the scope of the invention. Molecular weight values are determined employing get permeation chromatography (GPC) employing well-characterized polymethacytale (PMA) calibration standards:

Example 1

[0074] A container is charged with 35.1 parts methyl methacrylate, 136.5 parts Co-C11 methacrylate, 7.8 parts butyl methacrylate, 136.5 parts C12-C15 methacrylate, 70.2 parts C16-C18 methacrylate and 130 parts of a hydrotreated naphthenic oil (Risella G 07, Shell Germany) followed by stirring for 0.25 hours. A solution of 1.36 parts of 2,2-azobismethylbutyronitrile (VAZO-67, DuPont) in 2.7 parts toluene is added and stirred 0.1 hour. A reactor equipped with a stirrer, thermocouple reaching into the charged reaction mixture, No inlet atop an addition runnel attached to a subsurface tube, and H₂O condenser is charged with about 1/3 of the monomer-initiator solution. The remainder is placed in the addition funnel. With N2 addition at 0.3 standard cubic feet per hour (SCFH) and stirring, the mixture is heated to 100°C over 0.3 hour, heating is stopped and an exotherm to 131° C over 0.1 hours is observed. The temperature begins to drop and after 2 minutes, at 131°C dropwise addition of the remaining solution is begun. Addition time is 2 hours; 0.4 hours after the peak exothermic temperature, the temperature is 110° C. The temperature during addition is maintained at 110°C. The materials are cooled to 90°C over 0.3 hour followed by addition of 0.25 part t-butyl peroctoate (Trigonox 21, AKZO) followed by addition of 0.98 parts N-vinylimidazole (NVI), both all at once. While maintaining 90°C, three additional Increments, each of 0.98 parts NVI, are added at 0.25 hour Intervals for a total of 4 additions. The mixture is held at 90° C for 1,25 hours after the final NVI addition. An additional 0,13 parts Trigonox 21 Is added and the materials are held at 90°C for an additional 1 hour. An additional 80 parts Risella G 07 oil is added. the materials are heated to 150°C and stripped at 40-50 millimeters (mm) mercury for 1 hour, collecting 3 parts distillate. The residue is filtered employing a diatomaceous earth filter aid at 110°C. The resulting product has M.= 58.400 and

Example 2

polydispersity (M./M.) =2.90.

[0075] Following essentially the procedure of Example 1, a solution is prepared by mixing 27.5 parts methyl methacrylate, 5.5 parts buyly methacrylate, 105.5 parts C₁₂C₁₃ methacrylate, 52.5 parts C₁₂C₁₃ methacrylate and 80 parts Risella G 07 oil, then mixing in a solution of 1.2 parts VAZO-67 in 2.4 parts toluene. A reactor is charged with about 1/3 of this mixture and the remainder is added to an addition funnel. With stirring and

N₂ addition at 0.3 SCPH the mixture is heated to 100°C over 0.3 hours, heating is discontinued and the temperature rises exothermically to 139°C over 2 minutes. After 2 minutes the temperature begins to drop; dropwise addition of the remaining monomer-initiator mixture is begin. The temperature is at 110°C 0.3 hours after the peak exotherm. The balance of the mixture is added over 2 hours at 110°C. The materials are cooled to 90°C and 0.2 part Triggnox 21 is added all at one. To this mixture are added 3.5 parts NVI at at organise rate level 1 hour, the materials are heated at 90°C for 1 hour followed by addition of 0.1 part Triggnox 21. The materials are heated at 90°C for 2 hours followed by addition of 0.1 parts Triggnox 21 followed by stirring at 90°C for an additional 1.2 hours. To this mixture are added 81.5 parts Riscalla G 07 oil, the materials are heated with stirring to 150°C and stripped to 40-50 mm Hg for 1 hour while 2 parts distillate are collected. The residue is filtered using a diatomaceous earth filter aid. The filtrate has M_m-52,800 and polydiscensitive 2.75.

Example 3

[0078] A container is charged with 35.1 parts methyl methacrylate, 7.8 parts butyl methacrylate, 13.65, parts C₂C₁, methacrylate, 7.02 parts C₁₀C₂ methacrylate, 4.7 part NVI and 130 parts Riselia G 07 oil. The materials are stirred for 0.25 hour, then a solution of 1.56 part VAZO-67 in 3.12 parts toluren is added followed by stirring for 0.1 hour. A reactor equipped as described in Example 1 is charged with about 1/3 of this solution; the remainder is placed in the addition funnel. With stirring and V₂ addition at 0.3 SCPH, the mixture is heated to 110°C over 0.3 hour, heating is stopped and the temperature rises exotihermically to 13°C over 2 minutes. The temperature then begins to drop and after 2 minutes is a star 3°C. Dropwise addition of the remaining monomer-initiate mixture is begun and is continued for 2 hours. Temperature decreases to 110°C after 0.3 hours and is held at 110°C during addition. After addition at 6 continued for 2 hours. The materials are stirred at 90°C for 2 hours, 0.25 part Trigonox 21 is charged and the materials are heated for an additional 2 hours. The materials are ditted with 80 parts additional Risellia Gor 0.01, the add with stirring to 150°C, 40-55 mm Hg pressure, and stripped at 150°C for 1 hour, collecting 1 part distillate. The residue is filtered at 110°C with a distonmencous earth filter ald.

Example 4

[0077] A mixture of methacrylate ester monomers as described in Example 2, 80 parts of 150 Neutral mineral oil and 2.3 parts VAZC-67 in 15 parts butanol are combined to form a solution. Subsequent processing is substantially the same as described in Example 2 except the oil is 150 Neutral and the maximum exotherm is 136°C.

Example 5

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[0078] A container is charged with 30.5 parts methyl methacrylate, 117 parts Cg-C11 methacrylate, 6.1 parts butyl methacrylate, 117 parts C₁₂-C₁₅ methacrylate, 58.2 parts C₁₆-C₁₈ methacrylate and 88.7 parts of a hydrotreated naphthenic oil (Hydrocal 38, Calumet Lubricants, Princeton, LA, USA) followed by stirring for 0.25 hours. A solution of 1 part of 2,2-azobismethylbutyronitrile (VAZO-67, DuPont) In 5 parts toluene Is added and stirred 0.1 hour. A reactor equipped with a stirrer, thermocouple reaching into the charged reaction mixture, No inlet atop an addition funnel attached to a subsurface tube, and H₂O condenser is charged with about 1/3 of the monomer-initiator solution. The remainder is placed in the addition funnel. With N2 addition at 0.3 SCFH and stirring, the mixture is heated to 110°C over 0.3 hour, heating is stopped and an exotherm to 138° C over 0.1 hours is observed. The temperature begins to drop and after 3 minutes, at 136°C dropwise addition of the remaining solution is begun. Addition time is 2 hours; 0.4 hours after the peak exothermic temperature, the temperature is 110° C. The temperature during addition is maintained at 110°C. The materials are cooled to 90°C over 0.3 hour followed by addition of 0.2 part t-butyl peroctoate (Trigonox 21, AKZO) followed by dropwise addition of 3.9 parts NVI over 1 hour at 90°C. The reaction is held at 90°C for 1 hour. While maintaining 90°C, two additional increments of Trigonox 21, each of 0.1 part, are added, the second addition 2 hours after the first. The mixture is held at 90° C for 1 hour. An additional 90.4 parts Hydrocal 38 oll is added, the materials are heated to 150°C and stripped at 30-50 millimeters mercury for 1 hour, collecting 6 parts distillate. The residue is filtered employing a diatomaceous earth filter aid at 110°C. The resulting product has M_n= 71,600 and polydispersity =2.61.

Example 6

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[0079] A monomer-initiator solution is prepared as described in Example 1 except the oil is a 40 neutral naphthenic oil (Cross C-40, Cross Oil Co., Smackover AR, USA) and reacted substantially as described in Example 1; maximum exotherm = 138°C. After cooling to 90°C 0.25 part Trigonox 21 is added, stirred for 0.1 hour followed by dropwise

addition of 7 parts NVI over 1 hour at 90°C and heating for 1 additional hour. Trigonox 21 (0.13 part) is added, the materials are heated at 90°C for 2 hours then additional 0.13 part Trigonox 21 is added. The reaction is continued for 1 hour, 80 parts Cross L-40 oil are added and the materials are shipped to 150°C at 50 mm Hg for 1 hour removing 2 parts distillate. The materials are filtered yielding a filtrate containing a polymer having M_n =55,000 and polydispersity =2 22.

Example 7

[0080] The procedure of Example 6 is repeated except the oil is a hydrotreated naphthenic oil (Hydrocal 41, Calumet Lubricants). The product has M_p = 52,500 and polydispersity 2.73.

Example 8

[0081] Following essentially the procedure of Example 1, a solution is prepared by mixing 27.5 parts methyl methacyriate, 10.5 part about private, 10.5 parts (Q-G-m methacyriate, 10.5 parts (Q-G-m) methacyriate, 10.5 parts (Q-G-m) methacyriate, 10.5 parts (Q-G-m) methacyriate, 10.5 parts (Q-G-m) methacyriate and 80 parts (Q-G-m) ellowaters (Q-G-m) to 10 parts (Q-G

30 Example 9

[0082] A methacrylate ester-oil solution as described in Example 1 is prepared followed by addition of 1.55 parts VAZO-67 in 3.1 parts follower. Polymerization of the methacrylate monomers is conducted substantially as described in Example 1; peak exotherm = 135°C. After cooling to 90°C, 0.25 part Trigonox 21 and 1.75 part NVI are added followed by addition, at 0.25 hour intervals, of 3 additional increments of 1.75 parts NVI for a total of 7 parts. The reaction is continued for 1.25 hours, 0.15 part Trigonox 21 is added, reaction is continued for an additional 2 hours then 0.13 part Trigonox 21 is added and heating is continued for 1 hour all heating at 90°C. The materials are cooled, 80 parts filselia 6-07 oil are added and the materials are stripped to 150°C at 30.5 mm Hg for 1 hour collecting 1 part distillate followed by filtration through a distomaceous earth filter aid at 110°C. The filtrate contains a polymer having M₂ = \$5.00 and polydiepersity = 2.77.

Example 10

[0083] A container is charged with 33.9 parts methyl methacrylate, 7.5 parts butyl methacrylate, 133.6 parts Cg-C115 methacrylate, 135.6 parts Cg-C115 methacrylate, 135.6 parts Cg-C115 methacrylate, 135.6 parts N-vinyl pyrrolidnone and 130 parts Risella G0 70 ll. The materials are stirred for 0.5 brour, then a solution of 1.5 gear WAZO-67 in 3.1 parts toluene is added followed by stirring for 0.1 hour. A reactor equipped as described in Example 1 is charged with about 1/3 of this solution; the remainder is placed in the addition funnel. With stirring and N₂ addition at 0.3 SCFH, the mixture is heated to 110°C over 0.3 hour, heating is stopped and the temperature rises exothermically to 138°C over 3 minutes.

The temperature then begins to drop and after 2 minutes is at 136°C. Dropwise addition of the remaining monomer-intlator mixture is begun and is continued for 2 hours. Temperature decreases to 110°C after 0.3 hours and is held at 110°C during addition. After addition is completed, the mixture is cooled to 90°C over 0.3 hour followed by charging 0.25 part Tigronox 21: The materials are stirred at 90°C for 2 hours, 0.25 part Tigronox 21: The materials are heated for an additional 2 hours. The materials are stirred at 90°C tory stopped at 150°C, 40°C, and stripped at 150°C, 40°C on mml 4p pressure for 1 hour, collecting 1 part distillate. The residue is filtered at 110°C with a diatomaceous earth filter aid. The filtrate contains a polymer having M_n = 68,000 and M_w/M_n = 2.91

Example 11

[0084] The procedure of Example 10 is repeated employing 34.3 parts methyl methacrylate, 7.6 parts butyl methacrylate, 137 parts C₉₋₁₁ methacrylate, 135.3 parts C₁₆₋₁₈ methacrylate and replacing NVI with 8.73 parts N-vinyl formamide (NVF), other components remaining the same. The polymer product has M_m = 58,500 and M_m/M_m = 2.80.

Example 12

[0085] The procedure of Example 11 is repeated employing 34 parts methyl methacrylate, 7.5 parts butyl methacrylate, 13.9 parts each of C₈₋₁₁ and C₁₂₋₁₅ methacrylates, 67.9 parts C₁₆₋₁₆ methacrylate and replacing NVF with 12.9 parts 4-vinyl pyridine, other components remaining essentially the same. The polymer product has M_n = 56,500 and M_n/M_n = 2.64.

Example 13

[0086] The procedure of Example 11 is substantially followed replacing NVF with 17.3 parts N-vinyl caprolactam. The product has $M_n = 72,800$ and $M_w/M_n = 3.06$.

Example 14

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[0087] The procedure of Example 11 is substantially followed replacing NVF with 20.9 parts N,N-dimethylaminopropyl methacrylamide. The product has $M_n = 45,400$ and $M_w/M_n = 2.64$.

Example 15

[0088] The procedure of Example 14 is followed employing 10.5 parts N,N-dimethylaminopropyl methacrylamide. The product has $M_n = 45,200$ and $M_w/M_n = 2.66$.

Examples 16-21

[0089] Procedures similar to that of Example 11 are conducted replacing NVF with the indicated monomers:

Example	Monomer	Mn	M _w /M _n
16	acrylamide	79,300	3.04
17	t-octyl acrylamide	59,900	2.77
18	t-butyl acrylamide	53,100	2.85
19	methoxypropyl methacrylamide	57,800	2.8
20	2-(N,N-dimethylamino)ethyl methacrylate	62,300	2.86

Example 21

[0090] The procedure of Example 2 is repeated replacing hydrotreated naphthenic oil with 150 neutral mineral oil. The polymer has M_n = 56,800 and M_n/M_n = 2.57.

Example 22

[0091] The procedure of Example 2 is repeated replacing hydrotreated naphthenic oil with 85 neutral paraffinic oil (Total France). Polymer has M_n = 64,200 and M_m/M_n = 2.61.

Example 23

[0092] A container is charged with 57.5 parts of methyl methacytate, 12.2 parts butyl methacytate, 22.6.5 parts each of C₉₋₁₅ methacytate and C₁₂₋₁₅ methacytate, 11.4.8 parts C₁₆₋₁₆ methacytate, 11.7 parts NVI and 216.6 parts 85N parafilinic oil. The materials are stirred for 0.2 hours followed by addition of 2.27 parts 2,2²-azobis(2-methylbutyonlinic) (VAZO-67, DuPont) in 5 parts tolluene, and mixing 0.1 hour. One third of this mixture is charged to a reactor equipped with a stirrer, condenser. No intellect alone and difficult runnel and thermocouple in solution: the remainder is charced to

the addition furnel. With stirring and N₂ sparge at 0.3 SCPH the reactor contents are heated to 110°C over 0.5 hours, then heating is stopped. The temperature isses exchemically to 133°C over 0.05 hours. The temperature begins to drop and after 2 minutes reaches 132°C att which time addition of the solution from the addition funnel is begun. During addition (2 hours) the temperature drops to 110°C and is then maintained at 110°C. The reaction mixture is cooled to 90°C cover 0.3 hours then 0.43 patts, then 0.43 patts additional Trigonox 21 is charged, the materials are held 90°C for 2 hours, then 0.43 patts additional Trigonox 21 is charged, the materials are held at 90°C for 2 more hours. Additional 85N paraffinic oil is added (133.4 parts), the solution is heated to 150°C then strip to 80 mm Hg over 1 hour; 3 parts distillate collected. Product is filtered at 120°C with a distomacoous canth filter aid. M_s (PMA standard) 53,800 M_s/M_s, 2.8.

0 Example 24

[0093] A container is charged with 57.5 parts methyl methacrylate, 2.2 parts buth methacrylate, 2.26.5 parts each of C₉₋₁₁ methecrylate and C₁₉₋₂ methacrylate 1.4 [2.8] methacrylate 1.4 [2.9] methacrylate 1.4 [2.9] methacrylate 1.4 [2.9] parts tend-doded) metapata and 164.4 parts 8.5 neutral (85N) paraffinic oil followed by mixing at ambient temperature for 0.2 hour. To this solution are added 4.22 parts NAZO-67 dissolved in 9 parts tolusen followed by mixing for 0.1 hour. One hird (2.73 parts) of this solution is parts oil a reactor equipped with siterier, reflux condenser, thermocouple in solution and a N₂ inet along a addition funnel filled with the remaining 22 (547 parts) of methacrylate solution. To this solution in the reactor are added 11.7 parts N₂-(-dimethylamic)porpoylmethacrylamido, the materials are mixed for 0.2 hour then heated, under N₂ to 110°C over 0.5 hour whereupon heating is discontinued and the temperature rises exothermically to 141°C over 3 minutes. After the exothermic rise in temperature, the temperature is 140°C. The remaining materials are then added dropwise over 2 hours during which time the temperature drops to 110°C. After the addition is composed, the materials are held at 110°C for 0.5 hour, 0.43 part Trigonox 21 is added, the materials are held at 110°C for 1 hour, 0.43 part Trigonox 2 are added followed by heating at 110°C for an additional hour. The materials are heated to 125°C and stripped at 20 mm Hg for 0.5 hour, collecting 9 parts distillate followed by filtration at 120°C. The filtrate has M₁ = 38.6 and M₂ M₃ = 1.36.

[0094] The data in the following table illustrate the effect of the products of this invention with respect to improving viscosity bearesteristics of a typical tild full briefling viscosity. Each oil bland contains 5 % or 10 % by weight of the indicated product (no adjustment is made for diluent content of the additive in a solvent refined 100 neutral mineral oil. Viscosities are determined employing the procedure set out in ASTM Standard 0-445 and the viscosity index is determined employing the procedure set out in ASTM Standard 0-2270, both of which appear in the Annual Book of ASTM Standards, Section 5, Petrolaum Products. Luthicants and Fossifi Lusia, ASTM, 1916 Race Street, Philadelphia, PA, USA, ASTM Procedure 0-445 covers, in general, the determination of kinematic viscosity of flight petroleum products by measuring the time for a volume of liquid to five under gravity through a calibrated glass capillary visconnetic. ASTM Procedure D-2270 provides a means for calculating Viscosity Indox. Viscosities in the following table are given in entilistics:

Additive Effect-Viscosity and Viscosity Index

[0095]

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	@ 5 % Treatment				@ 10 % Treatment		
Product of		Visc	osity		Visco	osity	
Example	e 40°C 100°C		40°C 100°C Viscosity Index 40°C		100°C	Viscosity Index	
None	20	4.09	100				
1	30.4	6.5	176				
3	30.5	6.6	183				
4	28.8	5.9	158	40.8	8.4	185	
5	33.9	7.3	189	45.4	10.2	221	
6	31.6	6.8	180	47.5	10.3	213	
7	32.0	6.9	186	46.6	10.1	212	
8	28.7	6.0	164	40.5	8.6	199	
9	34.0	7.3	189	54.8	12.0	189	
10	32.3	7.0	189	48.4	10.8	222	
11	28.5	6.4	186	41.8	9.6	224	
13	30.6	6.6	178	47.5	10.4	216	

(continued)

	@ 5 % Treatment			@ 5 % Treatment @ 10 % Treatment		
Product of		Visc	osity		Visco	sity
Example	40°C	100°C	Viscosity Index	40°C	100°C	Viscosity Index
14	29.5	6.5	180	41.2	9.2	215
15	30.4	6.5	178	44.9	9.8	213
16	28.8	6.3	181	42.6	9.6	217
17	31.7	6.7	174	47.5	10.0	205
18	31.2	6.6	173	46.5	9.9	206
19	28.7	6.3	181			
20	33.3	7.1	185			
21	33.3	7.0	181	50.0	10.4	204
24	28.2	5.8	152	39.1	8.0	180

Low Temperature Characteristics

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[0096] As mentioned hereinabove, low temperature characteristics of lubricating compositions comprising additive concentrates of this invention which contain certain selected diluents are unexpectedly improved. This benefit is illustrated by the data in the following table.

[0097] Each blend contains 5% by weight of the product of the indicated Example (without adjusting for dilutent) in a partial example (without adjusting for dilutent) in a partial expense (as example (but a partial but a parti

Product of	oduct of Viscosity			Brookfield
Example	40°C	100°C	Viscosity Index	Viscosity (-40°C)
None	18.0	3.9	111	Solid
21	29.0	6.5	190	9250
(repeat)	29.4	6.6	192	12000
2	29.3	6.7	199	8450
(repeat)	29.5	6.8	200	10800
22	31.1	7.1	203	11400

Note: Repeat trials were conducted on freshly prepared blends; trials on blend containing product of Example 22 were conducted at the same time as repeat trials.

[0088] As is apparent, the polymers of these Examples are desirable viscosity index improvers. Another significant benefit is that the products of the instant invention improve both high temperature and low temperature characteristics of base oils. The viscosity of base oil without additive is strongly affected by changes in temperature while both the low and high temperature properties of the additive treated oils are less affected. In fact, the additives of this invention have a desirable viscosity lowering effect at very temperatures yet still fleduce the loss of viscosity at high temperatures. In effect, the additives perform not only as dispersant-viscosity improvers but also provide a fluidizing effect at very low temperatures.

[0099] A particularly surprising effect is provided by the product of Example 2.

[0100] In the following Table are given viscosity characteristics of diluents used to prepare the dispersant-viscosity improvers used in the lubricating oil compositions of the preceeding Table:

Oil	Used in Example	Viscosity 40°C	Viscosity 100°C	Viscosity Index	Brookfield 1 -26°C -	
150N	21	32.05	5.33	98	Channel	Solid
Risella G-07	2	2.91	1.18	-	90	150
85N	22	15.56	3.50	102	60,000	Solid

[0101] Noteworth is the fact that the typotrerated naphthenic oil has a viscosity index so low as to be vistored unmeasurable. Yet, then the viscosity improver of Example 2 is used in a lubricating oil composition, surprisingly the viscosity index of the resulting formulation is at least comparable to those incorporating dispersant viscosity improvers of Example 2 in and 22 which contain diluents having resideavely with viscosity indices.

6 [0102] Also surprising is that while the dispersant viscosity improvers of Examples 2 and 22 contain oils having relatively low kinematic viscosity, and provide good low temperature properties to a lubricating oil composition, there is no adverse effect on 40°C and 100°C viscosities.

The Oil of Lubricating Viscosity

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[0103] The lubricating compositions and methods of this invention employ an oil of lubricating viscosity, including natural or synthetic lubricating oils and mixtures thereof.

[0104] Natural oils include animal oils and vegetable oils (e.g. castor oil, lard oil) as well as mineral lubricating oils such as liquid pertoleum oils and solvent-treated or acid treated mineral lubricating oils of the peraffinic, naphthenic or mixed paraffinic-naphthenic types. Oils of lubricating viscosity derived from coal or shale are also useful. Synthetic lubricating oils include hydrocarbon oils and halosubstituted hydrocarbon oils such as polymerized and interpolymerized olefins, etc. and mixtures thereof, alitybenzenes, polyphenyl, (e.g. bjehnynk; tephenyls, alitybeted polyphenyls, etc.), silvylated diphenyl ethers and alitylated diphenyl sulfides and the derivatives, analogs and homologues thereof and the like.

20 [0105] Alkylene oxide polymers and interpolymers and derivatives thereof where their terminal hydroxyl groups have been modified by esterification, etherification, etc., constitute another useful class of known synthetic lubricating oils.
[0106] Another suitable class of synthetic lubricating oils that can be used comprises the esters of dI- and polycar-boxylic acids and those made from C₂ to C₂ monocerboxylic acids and polyolethers.

[0107] Other synthetic lubricating oils include liquid esters of phosphorus-containing acids, polymeric tetrahydrotrans and the lilet, silicon-based oils such as the polyalkyl-, polyaryl-, polyalkoxy-, or polyaryloxy-siloxane oils and silicate oils.

[0108] Unrefined, refined and rerefined oils, either natural or synthetic (as well as mixtures of two or more of any of these) of the type disclosed hereinabove can be used in the compositions of the present invention. Unrefined oils are similar to the outside directly from natural or synthetic sources without further purification treatment. Refined oils are similar to the unrefined oils except they have been further treated on one or more purification steps to improve one or more properties. Refined oils include solvent refined oils, hydrorefined oils, hydr

[0109] Rerefined oils are obtained by processes similar to those used to obtain refined oils applied to refined oils which have been already used in service. Such rerefined oils often are additionally processed by techniques directed to removal of spent additives and oil breakdown products.

[0110] Specific examples of the above-described oils of lubricating viscosity are given in Chamberlin, III, U.S. 4,326,972, European Patent Publication 107,282, and A. SequeriaJr., Lubricant Base Oil and Wax Processing, Chapter 6, Marcel Decker, Inc., New York (1994).

[0111] A basic, brief description of lubricant base oils appears in an article by D.V. Brock, "Lubrication Engineering", Volume 43, pages 184-5, March, 1987.

Other Additives

[0112] As mentioned, lubricating oil compositions of this invention may contain other components. The use of such additives is optional and the presence thereof in the compositions of this invention will depend on the particular use and level of performance required. Thus the other additive may be included or excluded. The compositions may comprise a zinc salt of a dithiophosphoric acid. Zinc salts of dithiophosphoric acids are often referred to as zinc dithiophosphates, zinc O,-diritydrocartly dithiophosphoric and the removal to provide additional extreme pressure, and intercommonly used names. They are sometimes referred to by the abbreviation ZDP. One or more zinc salts of dithiophosphoric acids may be present in a minor amount to provide additional extreme pressure, anti-wear and anti-violating verificance.

[0113] In addition to zinc salts of dithiophosphoric acids discussed hereinabove, other additives that may optionally be used in the lubricating oils of this invention include, for example, detergents, dispersants, viscosity improvers, oxidation inhibiting agents, pour point depressing agents, extreme pressure agents, anti-wear agents, color stabilizers and anti-foam agents. The above-mentioned dispersants and viscosity improvers may be used in addition to the additives of this invention.

[0114] Auxiliary extreme pressure agents and corrosion and oxidation inhibiting agents which may be included in the compositions of the invention are exempified by chlorinated aliphatic hydrocarbons, organic sulfides and onlysulfides, hosphorus esters including dihydrocarbon and trihydrocarbon nebsohites, mohydenum compounds. and the like

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[0115] Auxiliary viscosity improvers (also sometimes referred to as viscosity index improvers or viscosity modifiers) may be included in the compositions of this invention. Viscosity improvers are usually polymers, including polysobutions, polymethacytic acid esters, diene polymers, polyatkyl styrense, seterified styrene-maleic anhydricd copolymers, alkenylarene-conjugated diene-copolymers and polyotefins. Multifunctional viscosity improvers, other than those of the present invention, which also have dispersant and/or antioxidancy properties are known and may optionally be used in addition to the products of this invention. Such products are described in numerous publications including those mentioned in the Backcrown of the Invention.

[0116] Pour point cepressants are a particularly useful type of additive often included in the lubricating oils described herein. See for example, page 8 of 'Lubricant Additives' by C. V. Smalheer and R. Kennedy Smith (Lezius-Hiles Company Publisher, Cleveland, Ohio, 1967). Pour point depressants useful for the purpose of this invention, techniques for their preparation and their use are described in U. S. Patent numbers 2,387,501; 2,015,748; 2,655,479; 1,815,022; 2,191,498; 266,748; 2,721,877; 2,721,878; and 3,250,715.

[0117] Anti-foam agents used to reduce or prevent the formation of stable foam include silicones or organic polymers. Examples of these and additional anti-foam compositions are described in "Foam Control Agents", by Henry T. Kerner (Noyes Data Corporation, 1976), pages 125-162.

[0118] Detergents and dispersants may be of the ash-producing or ashless type. The ash-producing detergents are exemplified by oil solition reutral and basic salts of alkalin or alkaline earth metals with sullonic acids, carboxylic acids, phenois or organic phosphorus acids characterized by a least one direct carbon-to-phosphorus linkage.

[0119] The lerm "basic sait" is used to designate metal saits wherein the metal is present in stoichiometrically larger amounts than the organic acid radical. Basic salts and techniques for preparing and using them are well known to those skilled in the art and need not be discussed in detail here.

[0120] Ashless detergents and dispersants are so-called despite the fact that, depending on its constitution, the detergent or dispersant may upon combusation yield a nonvolatile residue such as boric valde or phosphonus pentoxide; however, it does not ordinarily contain metal and therefore does not yield a metal-containing ash on combustion. Many types are known in the art, and any of them are suitable for use in the lubricants of this invention. The following are illustrative:

(1) Reaction products of carboxylic acids (or derivatives thereof) containing at least 34 and preferably at least 54 center out the state of the carbox desired that carbo

i	3.163.603	3.381.022	3,542,680
ı	3.184.474	3,399,141	3.567.637
	3,215,707	3,415,750	3,574,101
	3,219,666	3,433,744	3,576,743
ı	3,271,310	3,444,170	3,630,904
	3,272,746	3,448,048	3,632,510
	3,281,357	3,448,049	3,632,511
	3,306,908	3,451,933	3,697,428
	3,311,558	3,454,607	3,725,441
	3,316,177	3,467,668	4,194,886
	3,340,281	3,501,405	4,234,435
	3,341,542	3,522,179	4,491,527
	3,346,493	3,541,012	RE 26,433
	3,351,552	3,541,678	

(2) Reaction products of relatively high molecular weight aliphatic or alicyclic halides with amines, preferably polyalkylene polyamines. These may be characterized as "amine dispersants" and examples thereof are described for example, in the following U.S. patents:

3,275,554	3,454,555
3,438,757	3,565,804

(3) Reaction products of alkyl phenois in which the alkyl groups contains at least 30 carbon atoms with aldehydes

(especially formaldehyde) and amines (especially polyalkylene polyamines), which may be characterized as "Mannich dispersants". The materials described in the following U. S. patents are illustrative:

3,413,347	3,725,480
3,697,574	3,726,882
3,725,277	l l

(4) Products obtained by post-treating the carboxylic amine or Mannich dispersants with such reagents as urea, thiourea, carbon disulfide, aldehydes, ketones, carboxylic acids, hydrocarbon-substituted succinic anhydrides, nitriles, epoxides, boron compounds, phosphorus compounds or the like. Exemplary materials of this kind are described in the following U.S. patents:

3,036,003	3,282,955	3,493,520	3,639,242
3,087,936	3,312,619	3,502,677	3,649,229
3,200,107	3,366,569	3,513,093	3,649,659
3,216,936	3,367,943	3,533,945	3,658,836
3,254,025	3,373,111	3,539,633	3,697,574
3,256,185	3,403,102	3,573,010	3,702,757
3,278,550	3,442,808	3,579,450	3,703,536
3,280,234	3,455,831	3,591,598	3,704,308
3,281,428	3,455,832	3,600,372	3,708,522
			4,234,435

(5) Polymers and copolymers of oil-solubilizing monomers such as decyl methacytak; viny' decyl ether and higher molecular weight olefins with monomers containing monopairs substituents, e.g., aminoality, deylates or methacytates, acrylamides and poly-(oxyethylene)-substituted acrylates. These may be characterized as "polymeric dispersants" and examples thereof are disclosed in the following U.S. patents:

3,329,658	3,666,730
3,449,250	3,687,849
3,519,565	3,702,300

[0121] The above-illustrated additives may each be present in lubricating compositions at a concentration of as little as 0.001% by weight, usually ranging from 0.01% to 20% by weight. In most instances, they each contribute from 0.1% to 10% by weight. The contribute from 0.1% to 10% by weight.

[0122] The various additives described herein can be added directly to the lubricant. Preferably, however, they are diluted with a substantially inert, normally liquid organic dilutent such as mineral oil, naphtha, benzene, toluen or xylene, to form an additive concentrate. Preferred additive concentrates usually comprise 0.1 to 80% by weight of the compositions of this invention and may contain, in addition, one or more other additives known in the art or described hereinabove. Concentrations such as 15%, 26%, 30% or 55% or higher may be employed.

[0123] The lubricating compositions of this invention are illustrated by the examples in the following Tables. The lubricating compositions are prepared by combining the specified ingredients, individually or from concentrates, in the indicated amounts and oil of lubricating viscosity to make the total 100 parts by weight. The amounts shown are parts by weight and, unless indicated otherwise, are amounts of chemical present on an oil-free basis. Thus, for example, an additive comprising 50% oil used at 10% by weight in a blend, provides 5% by weight of chemical. These examples are presented for illustrative purposes only, and are not intended to limit the scope of this invention.

Examples I-VI

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[0124] Lubricating oil compositions are prepared by blending into a mineral oil basestock 1.41% of the reaction product of a polysicoluteryl (M_P, a 1700) substituted succinic anhytidie and ethylene polyamine, 0.47% of sulfurzion product of a polysicoluteryl (M_P, a 1700) substituted succinic anhytidie and ethylene polyamine, 0.47% of sulfurzion bliefs-Alder adduct of butadiene and butyl acrylate, 0.81% of a zinc salt of mixed primary dialityl dithiophosphoric acids, 0.78% of calcium overbased (material raite 12) alityl progress sulforine scied and 3.3% of the listed products of this invention.

Example	Product of Example
1	3
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111	10
I۷	11
V	12
VI	16

Examples VII-X

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[0125] Lubricating oil compositions similar to those of the preceeding examples are prepared in a mineral oil basestock, adding thereto 1.41% of the reaction product of a polysobuteny (M_m = 1700) substituted succinic arrhydride and ethylene polyamine, 0.61% of suffurized Diles-Alder adduct of butadiene and buyl acrylate, 1.05 % of a zinc satt of mixed primary dialkyl dithiophosphoric acids, 1.0% of calcium overbased (metal ratio 12) alkyl benzene sulfonic acid and 3.3 % of the listed products of this invention:

Example	Product of Example	
VII	13	
VIII	14	
IX	15	
X	17	

[0126] While the invention has been explained in relation to its preferred embodiments, it is to be understood that various modifications thereof will become apparent to those skilled in the art upon reading the specification.

Claims

- A nitrogen-containing copolymer having a number average molecular weight in the range from 10,000 to 300,000 and polydispersity values in the range from 1.5 to 5 comprising units derived from
 - (A) 30% to 60% by weight of alkyl acrylate ester monomers containing from 1 to 11 carbon atoms in the alkyl group:
 - (B) 40% to 70% by weight of alkyl acrylate ester monomers containing from 12 to 24 carbon atoms in the alkyl croup; and
 - (C) 0.5% to 5% by weight of at least one nitrogen containing monomer selected from viny substituted nitrogen heterocyclic monomers, NN-disklyatimosklys arrylate monomers, NN-disklyatimosklys arrylatide monomers and tertiary-aiky a crylamides wherein (C) is incorporated as a comonomer with (A) and (B) or is grafted onto an skyl actyriate copolymer derived from (A) and (B) provided that the total equals 100%.
- 2. A copolymer of claim 1 wherein monomer (A) comprises from at least 5% by weight of alkyl acrylate esters having from 4 to 11 carbon atoms in the alkyl group.
 - A copolymer of claim 1 wherein monomer (A) comprises from 10% to 40% by weight of alkyl acrylate esters having from 1 to 4 carbon atoms in the alkyl group and from 60% to 90% by weight of alkyl acrylate esters having from 9 to 11 carbon atoms in the alkyl group.
 - 4. A copolymer of any preceding claim wherein the nitrogen containing monomer is selected from an N.N-diskylami-noskyl arraymide or arryished wherein each skilly or aminoskilly group contains, independently, from 1 to about 8 carbon atoms, tertiary butlyl acrylamide, vinyl prytidine, and an N-vinyl asubstituted heterocyclic monomer selected from N-vinyl acrylamide.
 - 5. A copolymer of any preceding claim wherein the alkyl acrylate ester monomers comprise alkyl methacrylate esters.

- A process for preparing a nitrogen containing copolymer having a number average molecular weight in the range from 10,000 to 300,000 and polydispersity values in the range from 1.5 to 5 comprising reacting in the presence of a free radical initiator
- 5 (A) from 30% to 60% by weight of alkyl acrylate ester monomers containing from 1 to 11 carbon atoms in the alkyl group.
 - (B) from 40% to 70% by weight of alkyl acrylate ester monomers containing from 12 to 24 carbon atoms in the alkyl group; and
 - (C) from 0.5% to 5% by weight of at least one nitrogen containing monomer selected from vinyl substituted nitrogen heterocyclic monomers, N,N-dialkylaminoalkyl acrylate monomers, N,N-dialkylaminoalkyl acrylamide monomers, N,N-dialkylaminoalkyl acrylamide monomers, N,N-dialkylaminoalkyl acrylamide monomers, N,N-dialkylaminoalkyl acrylamide, provided that the total equals 100%, optionalky in the presence of
- 7. A process for preparing a nitrogen containing copolymer having a number average molecular weight in the range from 10,000 1300,000 and polydispersity values in the range from 1.5 to 5 comprising grafting onto a polyacrylate copolymer comprising (A) from 30% to 65% by weight of units derived from alkyl acrylate ester monomers containing from 1 to 11 carbon atoms and (8) from 40% to 70% by weight of units derived from alkyl acrylate ester monomers containing from 12 to 24 carbon atoms, from 0.5% to 5% by weight, based on polyacrylate copolymer, of (C) a nitrogen containing monomer selected from vinyl substituted nitrogen heterocyclic monomers, N.N-dialkylaminoalkyl acrylamide monomers and tertiary alkyl acrylamides, provided that the total is 100%, said arristing conducted in the presence of a free middeal initiatior.
 - A process of either one of claims 6 and 7 wherein monomer (A) comprises at least 5% by weight of alkyl acrylate esters containing from 4 to 11 carbon atoms in the alkyl group.
 - 9. A process of any one of claims 6 to 8 conducted in an extruder.

a chain transfer agent.

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- 10. A process of claim 9 wherein the reaction is conducted in the absence of any diluent.
- 30 11. A process of claim 9 wherein the reaction takes place in a diluent.
 - 12. A process of claim 7 wherein the polyacrylate backbone has been prepared in the presence of an organic diluent.
 - 13. A process of claim 7 wherein the preparation of the polyacrylate backbone and the subsequent grafting are each conducted in an extruder.
 - 14. A process of claim 8 wherein monomers (A) and (B) and the fnee radical initiator are first combined to form a first mixture, whereupon from 20% to 40% of said first mixture is mixed with at least 50% of the Intended charge of monomer (C) to form a second mixture, healting said second mixture until an exotherm is noted whereupon after the exotherm subsides the materials are heated to maintain reaction temperature and the remaining first mixture of monomers (A), (B), and fullilator is added and the reaction is continued to completion.
 - 15. An additive concentrate comprising a copolymer of any one of claims 1 to 5 and a diluent.
- 45 16. An additive concentrate of claim 15 wherein the diluent is selected from mineral oils and synthetic oils and displays Brookfield viscosities measured by ASTM D-2983 ranging from 50 to 400 centipose at -26°C and from 100 to 1500 centipose at -40°C.
 - 17. An additive concentrate of claim 16 wherein the diluent is a mineral oil consisting essentially of hydrotreated naph-
 - 18. An additive concentrate of claim 16 wherein the diluent is a synthetic oil selected from esters, polyalphaolefin oligomers and alkylated benzenes.
- 19. An additive concentrate of any one of claims 15 to 18 comprising from 25% to 90% by weight of copolymer and from 10% to 75% by weight of diluent.
 - 20. A lubricating oil composition comprising a major amount of an oil of lubricating viscosity and a minor amount of a

nitrogen-containing copolymer of any one of claims 1 to 5.

A lubricating oil composition of claim 20 wherein the oil of lubricating viscosity comprises a mixture of mineral oil
and synthetic oils.

Patentansprüche

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- Ein Stickstoff-enthaltendes Copolymer mit einem Zahlenmittel des Molekulargewichts im Bereich von 10000 bis 300000 und Polydispersitätswerten im Bereich von 1.5 bis 5, umfassend Einheiten, die abgeleitet sind von:
 - (A) 30 bis 60 Gew.-% Alkylacrylatestermonomeren, die 1 bis 11 Kohlenstoffatome in der Alkylgruppe enthalten; (B) 40 bis 70 Gew.-% Alkylacrylatestermonomeren, die 12 bis 24 Kohlenstoffatome in der Alkylgruppe enthalten:
- (C) 0,5 bis 5 Gew.-% mindestens eines Stickstoff-enthaltenden Monomers ausgewählt aus vinylsubstitulerten Stickstoff-heterocyclischen Monomeren, NN-Dialikylaminoalkylacrylatmonomeren, N,N-Dialkylaminoalkylacrylamidmonomeren und tert-Alkylacrylamiden, wobei
 - (C) als Comonomer mit (A) und (B) einverleibt ist oder auf ein von (A) und (B) abgeleitetes Alkylacrylat-Copolymer aufgepfropft ist, mit der Maßgabe, dass insgesamt 100 % erhalten werden.
 - Copolymer nach Anspruch 1, wobel das Monomer (A) mindestens 5 Gew.-% Alkylacrylatester mit 4 bis 11 Kohlenstoffatomen in der Alkylaruppe umfasst.
- Copolymer nach Anspruch 1, wobei das Monomer (A) 10 bis 40 Gew.-% Alkylacrylatester mit 1 bis 4 Kohlenstoffatomen in der Alkylgruppe und 60 bis 90 Gew.-% Alkylacrylatester mit 9 bis 11 Kohlenstoffatomen in der Alkylgruppe umfasst.
 - 4. Copolymer nach einem der vorstehenden Ansprüche, wobei das Stücksteffenthattende Monomer ausgewährt ist aus einem N.-Dialtyleinminaeilylacryteinfül oder -acyteja, worn igde Altyl- oder -ninneistyleignyein per unbähräng! of bis etwa 8 Kohienstoffatome enthält, tert-Butylacryteinfül, Vinylpyridin und einem N-vinylaubstitulerten heterocyclischen Monomer ausgewährt aus N-Vinyliniadizol, N-Vinyleyerriolidinon und N-Vinylacprolatarin.
 - Copolymer nach einem der vorstehenden Ansprüche, wobei die Alkylacrylatestermonomere Alkylmethacrylatester umfassen.
 - Ein Verfahren zur Herstellung eines Stickstoff-enthaltenden Copolymers mit einem Zahlenmittel des Molekulargewichts im Bereich von 10000 bis 300000 und Polydispersitätswerten im Bereich von 1,5 bis 5, umfassend die Umsetzung von:
 - (A) 30 bis 60 Gew.-% Alkylacrylatestermonomeren, die 1 bis 11 Kohlenstoffatome in der Alkylgruppe enthalten; (B) 40 bis 70 Gew.-% Alkylacrylatestermonomeren, die 12 bis 24 Kohlenstoffatome in der Alkylgruppe enthal-
 - (C) 0,5 bis 5 Gew.-% mindestens eines Sickstoff-enhaltenden Monomers ausgewählt aus vinyisubstituierten Sickstoff-Retorcyclischen Monomeren, N.P-Diallydaminasilkydenrylatmoneneren, N.P-Diallydaminasilkylacrylamidmonomeren und tert-Alkykacrylamiden in Gegenwart eines Badikasitarters, mit der Maßgabe, dass die Summe 10% beträtig, gegebenenfalls in Gegenwart eines Keltenübertsgungsmittels.
 - 7. Ein Verfahren zur Herstellung eines Stickstoff-enthaltenden Copolymers mit einem Zahlenmittel des Molekulargewichts im Bereich von 1000 bis 30000 und Polydispensitätswerten im Bereich von 1,5 bis 5, umfassend das Pfropfen auf ein Polyacrylat-Copolymer, umfassend (A) 30 bis 50 Gew.-% Alkylacrylatestermonomere, die 1 bis 11 Kohlenstüffätome in der Alkylgruppe enthalten, 0,5 bis 5 Gew.-% auf köplacrylatestermonomere, die 12 bis 24 Kohlenstöffatome in der Alkylgruppe enthalten, 0,5 bis 5 Gew.-% auf der Basis des Polyacrylat-Copolymers (C) eines Stückstöff-enthaltendern Monomers ausgewählt aus virtyslubstüluerten Stückstöff-heterocyclischen Monomeren, N.Polisklyarimosalkylacrylationomeren, N.Polisklyarimosalky
 - 8. Verfahren nach Anspruch 6 oder 7, wobei das Monomer (A) mindestens 5 Gew.-% Alkylacrylatester mit 4 bis 11

Kohlenstoffatomen in der Alkylgruppe umfasst.

- 9. Verfahren nach einem der Ansprüche 6 bis 8, wobei das Verfahren in einem Extruder durchgeführt wird.
- Verfahren nach Anspruch 9, wobei die Umsetzung ohne Verdünnungsmittel durchgeführt wird.
 - 11. Verfahren nach Anspruch 9, wobei die Umsetzung in einem Verdünnungsmittel stattfindet.
 - Verlahren nach Anspruch 7, wobei das Polyacrylat-Grundgerüst in Gegenwart eines organischen Verdünnungsmittels heroestellt worden ist.
 - 13. Verfahren nach Anspruch 7, wobei die Herstellung des Polyacrylat-Grundgerüsts und das anschließende Pfropfen jeweils in einem Extruder durchgeführt werden.
- 13 14, Verfahren nach Anspruch 6, wobei die Monomere (A) und (B) und der Radikalstarter zunächst unter Bildung eines ersten Gemisches kombiniert werden, worauf 20 bis 40 % des ersten Gemisches mit mindestens 50 % der vorgesehnen Beschickung des Monomers (C) unter Bildung eines zweiten Gemisches gemischt werden, worauf das zweite Gemisch erhitzt wird, bis eine exotherme Reaktion beobachtet wird, worauf nach dem Abklingen der exothermen Reaktion die Materialten erhitzt werden, und ein Reaktionstemperatur aufrechtzuterhalten und azu restliche erste Gemisch der Monomeren (A) und (B) und der Starter zugegeben und die Umsetzung bis zur Vollständigkeit fordreseztzt wird.
 - 15. Ein Additivkonzentrat, das ein Copolymer nach einem der Ansprüche 1 bis 5 und ein Verdünnungsmittel umfasst.
- 25 16. Additivkonzentrat nach Anspruch 15, wobei das Verdünnungsmittel aus Mineralölen und synthetischen Ölen ausgewählt ist und eine gemäß ASTM D-2983 gemessene Brookfield-Viskosität von 0,05 bis 0,4 Pa · s (50 bis 400 centipoise) bei -26°C auf 0,1 bis 1,5 Pa · s (100 bis 1500 centipoise) bei -26°C aufweist.
- Additivkonzentrat nach Anspruch 16, wobel das Verdünnungsmittel ein Mineralöl ist, das im Wesentlichen aus mit
 Hydrotreating behandeltem nachthenischen Öl besteht.
 - Additivkonzentrat nach Anspruch 16, wobei das Verdünnungsmittel ein synthetisches Öl ist, das aus Estern, Polyalphaolefin-Oligomeren und alkyllerten Benzolen ausgewählt ist.
- 19. Additivkonzentrat nach einem der Ansprüche 15 bis 18, das 25 bis 90 Gew.-% Copolymer und 10 bis 75 Gew.-% Verdünnungsmittel umfasst.
 - Eine Schmierölzusammensetzung, die eine Hauptmenge eines Öls mit Schmierviskosität und eine geringere Menge eines Stickstoff-enthaltenden Copolymers nach einem der Ansprüche 1 bis 5 umfasst.
 - Schmierölzusammensetzung nach Anspruch 20, wobel das Öl mit Schmierviskosität ein Gemisch aus Mineralöl und synthetischen Ölen ist.

45 Revendications

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- Un copolymère renfermant de l'azote, ayant un polds moyen en nombre dans la gamme de 10 000 à 300 000 et des valeurs de de polydispersité dans la gamme de 1,5 à 5, comportant des motifs dérivés de :
- (A) 30 à 60 % en poids de monomère d'ester d'acrylate d'alkyle qui renferme de 1 à 11 atomes de carbone dans le groupe alkyle;
 - (B) de 40 % à 70 % en poids de monomère d'ester d'acrylate d'aikyle renfermant de 12 à 24 atomes de carbone dans le groupe alkyle;
 - (C) 0, 5 à 5 % en poids d'au moins un monomère renfermant de l'azote choisi parmi des monomères hétérocycliques azotés à substitution vinyle, des monomères de N-N dialkylaminolkyle acrylates, des monomères de N-N dialkylaminolkyle acrylamides et des tert-alkyl acrylamides, dans lequel (C) est incorporé en tant que monomère avec (A) et (B) ou bien est grefifé sur un copolymère d'acrylate d'alkyle dérivé de (A) et (B), sous la condition que le total soit écal à 100%.

- Un copolymère selon la revendication 1, dans lequel le monomère (A) comporte au moins 5 % en poids d'acrylate d'alkyle ayant de 4 à 11 atomes de carbone dans le groupe alkyle.
- 3. Un copolymère selon la revendication 1 dans lequel le monomère (A) comporte de 10 % à 40 % en poids d'esters d'acrylate d'airlyle ayant de 1 à 4 atomes de carbone dans le groupe alixyle et de 50 % à 90 % en poids d'esters d'acrylate d'alixyle avant de 9 à 11 atomes de carbone dans le groupe alixyle.
- 4. Un copolymère selon l'une quelconque des revendications précédentes, dans lequel le monomère renfemant de l'azote est choisi parmi un N-N-dialkylaminoalkyle acrylamide ou acrylate, dans lesquels chaque groupe alkyle ou aminoalkyle renferme, indépendamment, de 1 environ à 8 atomes de carbone, un tert-butyl acrylamide, de la virily pyridine et un monomère hétérocyclique à substitution N-vinyte, choisi parmi le N-vinyl imidazole, la N-vinyl pyrrolidone et la N-vinyl acronicatame.

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- Un copolymère selon l'une quelconque des revendications précédentes, dans lequel les monomères d'esters d'acrylate d'alkyle comportent des esters de méthacrylate d'alkyle.
 - Un procédé pour préparer un copolymère renfermant de l'azote, ayant un poids moléculaire moyen en nombre dans la gamme de 10 000 à 300 000 et des valeurs de polydispersité dans la gamme de 1,5 à 5, consistant à faire réagir, en présence d'un initiateur de radicaux libres,
 - (A) De 30 à 60 % en polds de monomères d'ester d'acrylate d'alkyle qui renferme de 1 à 11 atomes de carbone dans le groupe alkyle :
 - (B) de 40 % à 70 % en poids de monomères d'ester d'acrylate d'alkyle renfermant de 12 à 24 atomes de carbone dans le groupe alkyle :
 - (C) 0,8 à 5 % en polids d'au moins un monomère enfermant de l'azote choisi parmi des monomères héteroproliques azotes à substitution virvle, des monomères de N-N dissilvaiminolity les cerylates, des monomères de N-N dialitylaminolityle acrylamides et des tert-alityle acrylamides, sous la condition que le total soit égal à 100 %, facultativement en la présence d'un agent de transfert de chafine.
- 30 1. Un procédé pour préparer un copolymère renfermant de l'azote, ayant un poids moléculaire moyen en nombre dans la gamme de 10 00 à 300 00 et des valeurs de polydispersité dans la gamme de 1,5 à 5, consistant à grelfer sur un copolymère de polyacytale comportant (A) de 30 % à 60 % en poids de motils dérivés de monomères desters d'acytale callagie, entermant de 1 à 2 à 14 atomes de cachone et (B) de 40 % à 70 % en poids de motils dérivés de monomères d'ésters d'acytale d'altiyle, renfermant de 1 à 2 à 4 atomes de cachone, de 0,5 % à 5 % en poids en se basant sur un copolymère de polyacytale, de (C) un monomère néfermant de l'azote choisi parmi des monomères hétérocycliques azotés à substitution vinyle, les monomères de N-N-dialityalminoality explamides, des monomères de N-N-dialityalminoality explamides et des tert-altiyle acrylamides, sous la condition que le total soit de 100 % a lotting refle fétant fealisée en la présence d'un initiater ur de radieux libres.
- Un procédé selon une quelconque des revendications 6 et 7, dans lequel le monomère (A) comprend au moins 5
 en poids d'esters d'acrylate d'alkyle, qui renferme de 4 à 11 atomes de carbone dans le groupe alkyle.
 - 9. Un procédé selon l'une quelconque des revendications 6 à 8, mis en oeuvre dans une extrudeuse.
- 45 10. Un procédé selon la revendication 9, dans lequel la réaction est mise en oeuvre en l'absence d'un diluant quelconque.
 - 11. Un procédé selon la revendication 9, dans lequel la réaction se déroule dans un diluant.
- 50 12. Un procédé selon la revendication 7, dans lequel la structure polyacrylate a été préparée en présence d'un diluant organique.
 - 13. Un procédé selon la revendication 7, dans lequel la préparation de la structure polyacrylate et le greffage subséquent sont réalisés chacun dans une extrudeuse.
 - 14. Un procédé selon la revendications 6, dans lesquels les monomères (A) et (B) et l'initiateur de radicaux libres sont d'abord combinés pour former un premier métange, après quoi de 20 % à 40 % du dit premier métange sont métanges avec au moins 50 % d'une charge appropriée de monomères (C) pour former un second métange,

chaulfer le dit second mélange jusqu'à ce qu'un dégagement exothermique soit noté, après quoi, une fois que le dégagement exothermique d'iminue, les produits sont chauffés pour maintenir la température de la réaction et le premier mélange restant de monomères (A), (B), et un initiateur est ajouté et la réaction est poursuivie jusqu'à sa fin.

5 15. Un concentré additif comportant un copolymère selon l'une quelconque des revendications 1 à 5 et un diluant.

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- 16. Un concentré d'additif selon la revendication 15, dans lequel le diluant est choisi parmi les huiles minérales et les huiles synthétiques ti présente des viscosités Brookfield mesurées selon ASTM D-2983, se situant dans la gamme 50 à 400 centipoles à 2-60° ct et 10 à 1500 centipoises à -40°C.
- 17. Un concentré d'additif selon la revendication 16, dans lequel le diluant est une huile minérale constituée essentiellement d'une huile naphténique hydrotraitée.
- 18. Un concentré d'additif selon la revendication 16, dans lequel le diluant est une hulle synthétique choisie parmi les esters, des oligomères polyalphaoléfiniques et des benzènes alkyléniques.
- 19. Un concentré d'additif de l'une quelconque des revendications 15 à 18 comportant de 25 % à 90 % en poids de copolymères et de 10 % à 75 % en poids de diluant.
- 20. Une composition d'huile lubrifiante comportant une quantité prépondérante d'une huile de viscosité lubrifiante et une quantité plus faible d'un copolymère renfermant de l'azote selon l'une quelconque des revendications 1 à 5.
 - Une composition d'huile lubrifiante selon la revendication 20, dans laquelle l'huile de viscosité lubrifiante comprend un mélange d'huile minérale et d'huiles synthétiques.